

# UNITED STATES PARTMENT OF COMMERCE United States Patient and Trademark Office

Address: COMMISSIONER OF PATENTS AND TRADEMARKS

Washington, D.C. 20231

ATTORNEY DOCKET NO FIRST NAMED INVENTOR FILING DATE APPLICATION NO. 25436/1190 Ţ. SORGE 11/10/00 09/709,945 **EXAMINER** HM22/0620 SISSON, B KATHLEEN M WILLIAMS PH D PAPER NUMBER ART UNIT PALMER & DODGE LLF ONE BEACON STREET 1655 BOSTON MA 02108

DATE MAILED:

06/20/01

Please find below and/or attached an Office communication concerning this application or proceeding.

**Commissioner of Patents and Trademarks** 

|  | Application No.         | Applicant(s)   |
|--|-------------------------|--|
| Office Action Summary  | 09/709,945              | SORGE, JOSEPH A.   |
|  | Examiner                | Art Unit   |
|  | Bradley L. Sisson       | 1655   |
| The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply   |                         |  |
| A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) FROM THE MAILING DATE OF THIS COMMUNICATION.  - Extensions of time may be available under the provisions of 37 CFR 1.136 (a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication.  - If the period for reply specified above is less than thirty (30) days, a reply within the statutory minimum of thirty (30) days will be considered timely.  - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication.  - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133).  - Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b).  Status |                         |  |
| 1) Responsive to communication(s) filed on <u>07 M</u>   | <u> May 2001</u> .      |  |
| 2a) ☐ This action is <b>FINAL</b> . 2b) ☑ Th   | is action is non-final. |  |
| 3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.   |                         |  |
| Disposition of Claims  |                         |  |
| 4)⊠ Claim(s) <u>1-42</u> is/are pending in the application.  |                         |  |
| 4a) Of the above claim(s) <u>14-42</u> is/are withdrawn from consideration.  |                         |  |
| 5) Claim(s) is/are allowed.  |                         |  |
| 6) Claim(s) is/are rejected.   |                         |  |
| 7) Claim(s) is/are objected to.  |                         |  |
| 8) Claims are subject to restriction and/or election requirement.  |                         |  |
| Application Papers   |                         |  |
| 9) The specification is objected to by the Examiner.   |                         |  |
| 10) The drawing(s) filed on is/are objected to by the Examiner.  |                         |  |
| 11) ☐ The proposed drawing correction filed on is: a) ☐ approved b) ☐ disapproved.   |                         |  |
| 12) The oath or declaration is objected to by the Examiner.  |                         |  |
| Priority under 35 U.S.C. § 119   |                         |  |
| 13) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).  |                         |  |
| a) ☐ All b) ☐ Some * c) ☐ None of:   |                         |  |
| 1. Certified copies of the priority documents have been received.  |                         |  |
| 2. Certified copies of the priority documents have been received in Application No   |                         |  |
| 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).  |                         |  |
| * See the attached detailed Office action for a list of the certified copies not received.  14) Acknowledgement is made of a claim for domestic priority under 35 U.S.C. § 119(e).   |                         |  |
| 14) Acknowledgement is made of a claim for domestic phonty under 33 0.3.0. § 114(e).   |                         |  |
| Attachment(s)  |                         |  |
| <ul> <li>15) Notice of References Cited (PTO-892)</li> <li>16) Notice of Draftsperson's Patent Drawing Review (PTO-948)</li> <li>17) Information Disclosure Statement(s) (PTO-1449) Paper No(s)</li> </ul>   | 19) Notice of Informa   | ary (PTO-413) Paper No(s)<br>al Patent Application (PTO-152) |

Art Unit: 1655

#### **DETAILED ACTION**

### Election/Restrictions

- 1. Applicant's election of Group I, claims 1-13, in Paper No. 6 is acknowledged. Because applicant did not distinctly and specifically point out the supposed errors in the restriction requirement, the election has been treated as an election without traverse (MPEP § 818.03(a)).
- 2. Claims 14-42 have been withdrawn from further consideration pursuant to 37 CFR 1.142(b) as being drawn to a nonelected inventions, there being no allowable generic or linking claim. Election was made without traverse in Paper No. 6.

## Specification

3. The disclosure is objected to because it contains an embedded hyperlink and/or other form of browser-executable code. Applicant is required to delete the embedded hyperlink and/or other form of browser-executable code. See MPEP § 608.01.

## Claim Rejections - 35 USC § 112

4. The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

5. Claims 1-13 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to enable one skilled in the

Art Unit: 1655

art to which it pertains, or with which it is most nearly connected, to make and/or use the invention. As presently worded, the claims have sufficient breadth of scope to encompass arrays of oligonucleotides that can be of virtually any length, up to 599 nucleotides in length. The specification teaches at example 1, page 36, that an array had been manufactured, the number of different oligonucleotides synthesized and their lengths is not readily apparent. The aspect of one of skill in the art being able to effectively produce pure populations of oligonucleotides of lengths up to 599 nucleotides in length is critical to enabling the making and use of the claimed invention. At column 40 of Jones (US Patent 5,858,671) the inherent obstacle in synthesizing oligonucleotide arrays is disclosed. As stated therein, "that even if the constituent enzymatic steps approach 100% completion, incompletely processed products can accumulate to significant levels. For example, during oligonucleotide synthesis of a 70-mer, requiring 69 couplings, a 99% coupling efficiency results in only 50% of the generated oligonucleotides being full length  $(0.99^{69} = 0.50)$ ." In the present case, applicant is claiming a product that would be the result of 598 couplings, not just 69 as described above.

At page 36 of the specification it is asserted that "[t]he human cDNA microarray [was] produced from cloned selected at random from the clone collection, as diagramed in Figure 1A. Plasmid DNA of each clone is isolated by means known in the art." At page 40 it is stated that "[t]he 3" cDNA PCR products (nucleic acid members) [were] stably associated with a substrate which is a standard 25 mm X 75 mm glass microscope slide either by an arrayer or manually...." Such teachings, however, especially in light of the art-recognized "inherent obstacle in synthesizing oligonucleotide arrays," are not sufficient to enable the production of the now claimed array. While the specification need not set forth each and every possible combination of method steps

Page 4

Application/Control Number: 09/709,945

Art Unit: 1655

that may be utilized, the specification does need to provide sufficient detail so that the claimed method does not unfairly force the public to engage in undue experimentation. In the present case, the state of the prior art has advanced to the stage where certain problems are recognized as being endemic. The specification does not address how these art-recognized problems are to be overcome. Accordingly, the burden of enablement is unfairly shifted to that of the public. The situation at hand is analogous to that in *Genentech v. Novo Nordisk A/S* 42 USPQ2d 1001. As set forth in the decision of the Court:

"'[T]o be enabling, the specification of a patent must teach those skilled in the art how to make and use the full scope of the claimed invention without undue experimentation.' In re Wright 999 F.2d 1557, 1561, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993); see also Amgen Inc. v. Chugai Pharms. Co., 927 F. 2d 1200, 1212, 18 USPQ2d 1016, 1026 (Fed Cir. 1991); In re Fisher, 427 F. 2d 833, 166 USPQ 18, 24 (CCPA 1970) ('[T]he scope of the claims must bear a reasonable correlation to the scope of enablement provided by the specification to persons of ordinary skill in the art.').

\*\*\*\*

"Patent protection is granted in return for an enabling disclosure of an invention, not for vague intimations of general ideas that may or may not be workable. See Brenner v. Manson, 383 U.S. 519, 536, 148 USPQ 689, 696 (1966) (starting, in context of the utility requirement, that 'a patent is not a hunting license. It is not a reward for the search, but compensation for its successful conclusion.') Tossing out the mere germ of an idea does not constitute enabling disclosure. While every aspect of a generic claim certainly need not have been carried out by an inventor, or exemplified in the specification, reasonable detail must be provided in order to enable members of the public to understand and carry out the invention. "It is true . . . that a specification need not disclose what is well known in the art. See, e.g., Hybritech, Inc. v. Monoclonal Antibodies, Inc., 802 F.2d 1367, 1385, 231 USPQ 81, 94 (Fed. Cir. 1986). However, that general, oft-repeated statement is merely a rule of supplementation, not a substitute for a basic enabling disclosure. It means that the omission of minor details does not cause a specification to fail to meet the enablement requirement. However, when there is no disclosure of any specific starting material or any of the conditions under which a process can be carried out, undue experimentation is required; there is a failure to meet the enablement requirement that cannot be rectified by asserting that all the disclosure related to the process is within the skill of the art. It is the

Art Unit: 1655

specification, not the knowledge of one skill in the art, that must supply the novel aspects of an invention in order to constitute adequate enablement. This specification provides only a starting point, a direction for further research. (emphasis added)

For the above reasons, and in the absence of convincing evidence to the contrary, the specification has not been found to enable the making and use of the claimed device.

6. Claims 1-13 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention. As presently worded, the claimed array can be comprised of nucleic acid sequences that are less than 600 bases in length and which are not present in any public database (claims 1 and 8). In view of an independent claim encompassing all of the limitations of any of its dependent claim, claims 1-13 have been interpreted as encompassing non-known/not publicly available sequences.

As set forth in claim 7, the array can comprise from 1,000 to 10,000 positions/oligonucleotides. Claims 1-6 and 8-13 place no upper limit on the number of oligonucleotides present. Upon review of the specification no description of said non-known sequences is provided. Accordingly, the specification does not reasonably suggest that applicant was in possession of said non-publicly known sequences at the time of filing. In support of this position, attention is directed to the decision of *Vas-Cath Inc. v. Mahurkar* 19 USPQ2d 1111 (CAFC, 1991):

This court in *Wilder* (and the CCPA before it) clearly recognized, and we hereby reaffirm, that 35 USC 112, first paragraph, requires a "written description of the

Art Unit: 1655

invention" which is separate and distinct from the enablement requirement. The purpose of the "written description" requirement is broader than to merely explain how to "make and use"; the "applicant must also convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the "written description" inquiry, whatever is now claimed.

For the above reasons, and in the absence of convincing evidence to the contrary, the specification fails to reasonably suggest that applicant was in possession, at the time of filing, an array that comprised either in its entirety, or in part, oligonucleotide sequences that are not known to the public.

- 7. The following is a quotation of the second paragraph of 35 U.S.C. 112:
  The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.
- 8. Claims 4 is rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Claim 4 is indefinite with respect to what constitutes the metes and bounds of "substantially noncoding sequences" as a sequence either encodes a protein or it does not.

#### Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Bradley L. Sisson whose telephone number is (703) 308-3978. The examiner can normally be reached on 6:30 a.m. to 5 p.m., Monday through Thursday.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Stephanie Zitomer can be reached on (703) 308-3985. The fax phone numbers for

Art Unit: 1655

the organization where this application or proceeding is assigned are (703) 305-3592 for regular communications and (703) 308-0294 for After Final communications.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is (703) 308-0196.

Bradley L. Sisson Primary Examiner

Art Unit 1655

BLS

June 17, 2001